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(54) Title: CHONDROITIN LYASE ENZYMES

## (57) Abstract

The present invention describes a method for the production of two highly purified enzymes capable of degrading chondroitin sulfate polysaccharides. A multi-step purification method incorporating cell disruption, cation exchange chromatography, affinity chromatography, hydroxylapatite chromatography, high resolution ion exchange chromatography and size exclusion is outlined. A  $77,000 \pm 5,000$  Dalton protein capable of degrading chondroitin sulfates A and C and a  $55,000 \pm 2,300$  Dalton protein capable of degrading dermatan sulfate were isolated. The genes encoding these enzymes, chondroitinase AC and chondroitinase B, respectively, have been cloned and methods for their use are described.

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#### CHONDROITIN LYASE ENZYMES

## Background of the Invention

The present invention is the purification and cloning of chondroitin lyase enzymes found in Flavobacterium heparinum.

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Glycosaminoglycans are unbranched polysaccharides consisting of alternating hexosamine and hexuronic residues which carry sulfate groups in different positions. of molecules can be divided into three families according to the composition of the disaccharide These are: heparin/heparan sulfate backbone. [HexA-GlcNAc(SO<sub>4</sub>)]; chondroitin sulfate [HexA-GalNAc]; and keratan sulfate [Gal-GlcNAc]. chondroitin sulfate family includes seven sub-types designated unsulfated chondroitin sulfate, oversulfated chondroitin sulfate and chondroitin sulfates A-E which vary in the number and position of their sulfate functional groups. Additionally, chondroitin sulfate B, also referred to as dermatan sulfate, differs in that iduronic acid is the predominant residue in the alternative hexuronic acid position.

Chondroitin sulfates A, B and C are the 25 predominant forms found in mammals and may be involved in the modulation of various biological activities including cell differentiation, adhesion, enzymatic pathways and hormone interactions. The presence of chondroitin sulfate proteoglycans is elevated in the later stages of 30 cell growth in response to tissue and vessel damage, as reported by Yeo, et al., Am. J. Pathol. 138:1437-1450, 1991, Richardson and Hatton, Exp. Mol. Pathol. 58:77-95, 1993 and Forrester, et al., J. Am. Coll. Cardiol. 17:758-769, 1991. 35 Chondroitin sulfates also have been associated with events involved in the progression of vascular disease and lipoprotein uptake as described by

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Tabas, et al., J. Biol. Chem., 268(27):20419-20432, 1993.

Chondroitin enzymes of a suitable purity and characterization could be useful tools in determining the role of chondroitin sulfates in modulating these cellular events and in developing therapeutics for the treatment of disease states.

Chondroitin sulfate degrading enzymes,

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referred to as chondroitinases or chondroitin sulfate lyases, from several bacterial species have been reported. Takegawa, et al., J. Ferm. Bioeng. 77(2):128-131, 1991, report a chondroitinase AC from Aureobacterium with a molecular weight of between 81,000 and 83,000 Daltons that is inhibited by copper ions. Bacteriodes thetaiotamicron produces two chondroitinase AC degrading enzymes of molecular weight 104,000 and 108,000 Daltons, as described by Linn, et al., J. Bacteriol. 165:859-866, 1985. Other bacterium including

- 20 Flavobacterium heparinum, Proteus vulgaris,
  Arthrobacter aurescens and Pseudomonas fluorescens
  produce chondroitinase AC or chondroitinase ABC
  enzymes which are not well characterized, as
  reviewed by Linhardt, et al., Appl. Biochem.
- 25 Biotechnol. 12:135-177, 1986. F. heparinum is the only microbe that produces an enzyme which is specific for dermatan sulfate, chondroitinase B, as reported by Linhardt, R., et al. However, the chondroitinase degrading enzymes from F. heparinum have not been purified to homogeneity or thoroughly characterized.

It is therefore an object of the present invention to provide methods for purifying chondroitin lyase enzymes.

It is a further object of the present invention to provide DNA sequences encoding chondroitin lyase enzymes.

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It is a still further object of the present invention to provide purified chondroitin lyase enzymes which are useful as pharmaceutical regents.

## Summary of the Invention

A method for purifying chondroitin lyase enzymes from bacteria such as the Gram negative organism, Flavobacterium heparinum, have been developed which yields purified chondroitinase AC and chondroitinase B. Cells are grown by fermentation culture, the cells are lysed preferably using an osmotic shock technique which selectively releases proteins from the periplasmic space, then fractionated by cation exchange chromatography. Fractions containing chondroitinase degrading activity are further fractionated by affinity chromatography using a sulfated cellulose based resin and hydroxylapatite chromatography which separate the chondroitinase AC and chondroitinase B activities. Highly purified preparations of each enzyme are obtained by an additional chromatography step using a high resolution strong cation exchange resin. Pure preparations of chondroitinase B may require an additional separation step based on molecular size, such as gel filtration liquid chromatography.

The genes encoding chondroitinase AC and chondroitinase B enzymes of Flavobacterial origin were cloned. These can be used in conjunction with suitable expression systems to produce the enzymes in Flavobacterium, for example, under the control of overexpression promoters, or in organisms other than Flavobacterium.

## Brief Description of the Drawings

Figure 1 is a schematic of the construction of plasmids used to sequence the chondroitinase AC

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gene from Flavobacterium heparinum, pA2C1B, p64BS2-7. Restriction sites are: S - SaU, B - BamHI, P - PstI, E - EcoRI, H - HindIII, C - ClaI and K - KpnI.

Figure 2 is a schematic of the construction of pGB-ChAC, a plasmid capable of directing the expression of active chondroitinase AC in *E. coli* from tandem tac promoters (double arrowheads).

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Figure 3 is a schematic of the construction of plasmids used to sequence the chondroitinase B gene from Flavobacterium heparinum, pCHB300 and pCHB78.

Figure 4 is a schematic of the construction of pGB-CHB, a plasmid capable of directing the expression of active chondroitinase B in E. coli from tandem tac promoters (double arrowheads).

## Detailed Description of the Invention

# Purification of Chondroitin Sulfate degrading Enzymes from F. heparinum

20 Cells are grown in fermentation cultures to obtain sufficient quantities of the enzymes. Chondroitin sulfate A is included in the media at a concentration of between 0.5 and 10 g/l, preferably between 1.0 g/L to 2.0 g/l to induce chondroitinase AC and chondroitinase B synthesis. Crude enzyme 25 extracts are prepared by liberating soluble proteins from the cells by standard cell disruption techniques, preferably osmotic shock based techniques which selectively release proteins from the cell's periplasmic space. For example, 30 proteins can be released from the periplasmic space by treatment with non-ionic detergents in the range of 0.01 to 1.0%, freezing and thawing the cells, partial sonication for 0.5 to 6.0 minutes at 30 to 60% power in a pulsed mode 25/75 to 75/25, lysosyme 35

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treatment at 0.001 to 1.0 mg/ml for 15 to 60 minutes between 4 and 25°C, organic solvent treatment with 0.01 to 1.0% chloroform or toluene or by the osmotic shock process described in U.S. Patent No. 5,169,772 to Zimmermann and Cooney. In the latter, cells are partially sonicated for between 0.5 and 4.0 minutes, poser 3-6 pulsed mode 50/50, partial homogenization 250 to 500 psi, followed by lysozyme treatment at 0.001 to 1.0 mg/ml for between 15 and 60 minutes at between 4 and 23°C, and organic solvent treatment with 0.01 to 1.0% chloroform or 0.01 to 1.0% toluene.

In the preferred embodiment, the crude extract is fractionated by cation exchange chromatography using a high flow rate resin such as Sepharose™ S Big Beads (Pharmacia), MonoS™ (Pharmacia), CBX (J.T. Baker), Sepharose™ S (Pharmacia), and CM cellulose (Bio-Rad or Sigma), at a pH of between 6.0 and 8.5 with a salt gradient equivalent to 0.01 to 1.0 M NaCl. The bound proteins are preferably eluted with step gradients of 0.25 M sodium chloride and 1.0 M sodium chloride, at pH 7.0. Chondroitinase activity elutes in the 0.25 M sodium chloride fraction. Other salts can be utilized, such as sodium phosphate or sodium sulfate to create the salt gradient. Alternatively, a pH gradient in the range of 6.0 to 10.0 could be employed or a combination of a salt and pH gradient.

Fractions containing chondroitinase degrading activity are further fractionated by affinity chromatography using a sulfated cellulose based resin with a linear gradient of 0.0 to 0.4 M NaCl. Chondroitinase AC primarily elutes at 0.23 to 0.26 M NaCl and chondroitinase B elutes at 0.27 to 0.3 M NaCl. This is followed by hydroxylapatite chromatography using a step gradient of 0.25 M NaCl

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followed by a linear gradient of 0.25 to 1.0 M NaCl at pH 7.7. Chondroitinase B elutes at 0.25 M NaCl while chondroitinase AC elutes at 0.85 to 0.95 M NaCl. Highly purified preparations of each enzyme are obtained using a high resolution strong cation 5 exchange resin eluted with a linear gradient from 0.125 to 0.325 M NaCl in 0.025 M sodium phosphate at pH 7.0  $\pm$  0.1, as described with reference to elution from cation exchange resins described 10 above. Chondroitinase B elutes in a protein peak at 0.175 to 0.225 M NaCl. Chondroitinase B can be further purified on the basis of molecular size by size exclusion chromatography, ultrafiltration or preparative gel electrophoresis. Gel filtration 15 (size exclusion) resins with maximum resolution performance in the range of 5,000 to 100,000 are preferred. These include Superose™ 12, Superose™ 6, Sephadex<sup>™</sup> G-50 and Sephadex<sup>™</sup> G-50 from Pharmacia and BioGel™ P-60 and BioGel™ P-100 from BioRad. Ultrafiltration or dialysis membranes with 20 molecular weight cutoffs in the range of 10,000 to 30 000 Daltons are useful in removing small contaminants while ultrafiltration and dialysis membranes with molecular weight cut-offs in the 25 range of 70,000 to 1,000,000 Daltons are useful to remove larger contaminants. Alternatively, chondroitinase B containing samples of sufficient purity, more than 25% pure, could be further purified by subjecting the sample to gel electrophoresis according to standard laboratory 30 procedures, and excising the major band appearing at a molecular weight of 55,000  $\pm$  2,300 Daltons. The method of producing and purifying the chondroitinase lyase enzymes is exemplified as

follows.

F. heparinum was cultured in 15 L computer controlled fermenters in a variation of the defined

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nutrient medium described by Galliher, et al., Appl. Environ. Microbiol. 41(2):360-365, 1981. Chondroitin sulfate A (Sigma) was included in the media at a concentration of 1.0 g/L as the inducer of chondroitinase AC and chondroitinase B synthesis. The cells were harvested by centrifugation and the desired enzymes released from the periplasmic space by a variation of the osmotic shock procedure described by U.S. Patent No. 5, 169,772 to Zimmermann and Cooney. Cells 10 were resuspended in 0.01 M sodium phosphate and 0.3 M sodium chloride at pH 7.0  $\pm$  0.1 to give a final cell concentration of 100 absorbance units at 600 The non-ionic detergent Nonedit™ P-40 was added to the cell suspension to a final 15 concentration of 0.1% and the cells stirred for 1 hour at room temperature using a magnetic stir bar device. Cells and cell debris were then removed by centrifugation using a Sorval™ RC5C centrifuge with a JA-10 rotor at 10,000 RPM for 45 minutes. 20 cell pellet was discarded and the osmolate supernatant retained for further processing.

Osmolates obtained from F. heparinum fermentations induced with chondroitin sulfate A were subjected to centrifugation to remove cells and cell debris and the supernatant applied to a cation exchange column (5.0 cm x 30 cm, Sepharose<sup>TM</sup> S Big Beads, Pharmacia) at a linear flow rate of 10 cm@min^1. The bound proteins were eluted at a linear flow rate of 5.1 cm@min^1 with step gradients of 0.01 M phosphate, 0.01 M phosphate/0.25 M sodium chloride and 0.01 M phosphate/1.0 M sodium chloride, all at pH 7.0 ± 0.1. Chondroitinase activity eluted in the 0.25 M sodium chloride fraction.

This fraction was further purified by diluting the chondroitinase containing fraction

two-fold with 0.01 M sodium phosphate and applying the material onto a column containing cellufine sulfate (2.6 cm i.d. x 100 cm, Amicon) and eluting at a linear flow rate of 1.88 cm•min<sup>-1</sup> with a linear gradient of sodium chloride, 0.0 to 0.4 M. Chondroitinase AC primarily eluted at 0.23 to 0.26 M sodium chloride while chondroitinase B eluted at 0.27 to 0.3 M sodium chloride.

Each fraction was diluted two-fold with

0.01 M sodium phosphate and applied to a
hydroxylapatite column (2.6 cm i.d. x 30 cm). The
bound proteins were eluted with a step gradient of
0.25 M sodium chloride followed by a linear
gradient of 0.25 to 1.0 M sodium chloride all in

0.025 M sodium phosphate at pH 7.7 ± 0.1.
Chondroitinase B elutes in the 0.25 M sodium
chloride step while chondroitinase AC elutes at
0.85 to 0.95 M sodium chloride.

The chondroitinase B fraction was diluted two-fold in 0.01 M sodium phosphate and applied to 20 a strong cation exchange column (CBX-S, J.T. Baker, 1.6 cm i.d. x 10 cm). The bound material was eluted at a flow rate of 1.0 cmemin with a linear gradient from 0.125 to 0.325 M sodium chloride in 0.025 M sodium phosphate at pH 7.0  $\pm$  0.1. 25 Chondroitinase B eluted in a protein peak at 0.175 to 0.225 M sodium chloride and contained a minor contaminating protein of molecular weight 20,000 Daltons. This protein was removed by gel filtration chromatography by loading the 30 chondroitinase B sample onto a Superdex™ 200 column (1.0 cm i.d. x 30 cm, Pharmacia) and eluting with 0.05 M sodium phosphate, pH 7.2 at a linear flow rate of 1.25 cm min and collecting the protein containing fractions. 35

The chondroitinase AC fraction collected from hydroxylapatite chromatography was diluted

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three-fold in 0.01 M sodium phosphate and applied to a strong cation exchange column (CBX-S, J.T. Baker, 1.6 cm i.d. x 10 cm). The bound material was eluted at a flow rate of 1.0 cm min<sup>-1</sup> with a linear gradient from 0.125 to 0.325 M sodium chloride in 0.025 M sodium phosphate at pH 7.0 ± 0.1. Chondroitinase AC eluted in a single protein peak at 0.175 to 0.225 M sodium chloride. Purification results for the chondroitinase enzymes are shown in Table 1.

Table 1: Purification of chondroitinase enzymes from Flavobacterium heparinum fermentations

sample		rity		fic activ		yield (%)
<u>fermentatio</u>						
chondroitin				0.764		100
chondroitin	ase B	21,53	31	0.252		100
osmolate:						
chondroitin	ase AC	39.46	58	1:44		60
chondroitin				0.588		71
CHORGEOTETH	abc b .	10,20		0.500		
cation exch	ange:					
chondroitin	ase AC	27,93	35	9.58		43
chondroiting				4.731		64
Ollollar Oz Czil		,	-			
cellufine s	ulfate:	<u>.</u>				
chondroitin			50	22.6		28
chondroitin				21.2		<b>2</b> 9
CHONGLOTCIN		0,2	•			
hydroxylapa	tite:					
chondroitin		14,49	94	146.8		22
chondroitin				65.62		18
CHOMATOTOTA		5,5		••••	•	
strong cati	on exch	ange :	<u>.</u>			
chondroitin	ase AC	9,84	13	211.4	•	15
chondroiting				167.2		18
		-,				
gel filtrat	ion:					•
chondroiting		2,814	1	278.7		13
CHONGLOTCHI	450	2,01.	-	,		

Chondroitinase activity was determined by a modification of the spectrophotometric assay described by Yang, et al., *J. Biol. Chem.*, 160(30):1849-1857, 1985. Chondroitinases degrade

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their respective substrates by an eliminative reaction resulting in the formation of 4,5unsaturated sulfated disaccharides which absorb ultraviolet light at 232 nm. Reaction buffers contained 50 mM Tris, pH 8.0 and 0.5 mg/ml substrate; dermatan sulfate for chondroitinase B activity, chondroitin sulfate A for chondroitinase AC activity. A continuous spectrophotometric assay is carried out by transferring a 10 to 50  $\mu$ l sample to a quartz cuvette and adding the reaction buffer to make a final volume of one ml. The cuvette is placed in a Beckman DU 640 spectrophotometer, controlled to maintain a constant temperature of 30°C, and the increase in absorbance at 232 nm monitored for three to five minutes. Activities are calculated using the molar extinction coefficient for chondroitin sulfate, 5.1 x 103 M-1, and are expressed in international units, IU, where one IU is the amount of enzyme required to catalyze the formation of one  $\mu$ mole unsaturated product per minute.

#### Properties of Chondroitinase Enzymes

The purification method described herein is suitable for obtaining sufficient quantities of purified chondroitinase AC and chondroitinase B for characterization studies. The purified enzymes were analyzed by SDS-PAGE using the technique of Laemmli, Nature, 227:680-685, 1970, and the gels quantified with a scanning densitometer (Bio-Rad, Model GS-670). Chondroitinase AC was shown to have a molecular weight of  $77,000 \pm 5,000$  Daltons and a purity of greater than 99% while chondroitinase B has a molecular weight of  $55,000 \pm 2,300$  Daltons and a purity of greater than 99%.

Kinetic parameters of the 77,000 Dalton chondroitinase AC protein were measured using both chondroitin sulfate A and chondroitin sulfate C as

substrates. The  $K_m$  and  $K_{cat}$  values for chondroitinase A activity were 6  $\mu M$  and 230 s<sup>-1</sup>, respectively, while the  $K_m$  and  $K_{cat}$  values for chondroitinase C activity were 9.3  $\mu M$  and 150 s<sup>-1</sup>, respectively. Kinetic parameters of the 55,000 Dalton chondroitinase B protein were measured using dermatan sulfate as the substrate. The  $K_m$  and  $K_{cat}$  values for chondroitinase B activity were 7.4  $\mu M$  and 192 s<sup>-1</sup>, respectively.

### Effect of Added Reagents

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The  $V_{\text{max}}$  of the chondroitinase enzymes can be effected by trace amounts of certain elements. A base reaction buffer of 20 mM Tris buffer, pH 8.0 and 0.5 mg/ml substrate, either chondroitin sulfate A for chondroitinase AC or dermatan sulfate for chondroitinase B, was used to determine the effect of divalent metals and salts on the activity of the chondroitinase enzymes. The results are shown in Table 2.

Table 2: Effects of 0.1 mM of various reagents on the activity of chondroitinase enzymes.

reagent	chondroitinase AC relative activity(%)	chondroitinase B relative activity(%)
none	100	100
MgCl,	91	91
MnCl <sub>2</sub>	83	33
CuSO,	92	91
ZnCl,	26	45
FeSO <sub>4</sub>	98	69
HgCl,	55	40
CoCl <sub>2</sub>	81	42
EDTA	97	1

#### Stabilization of Chondroitinases

The chondroitinase enzyme activity can be stabilized by addition of excipients or by lyophilization. Stabilizers include carbohydrates, amino acids, fatty acids, and surfactants and are known to those skilled in the art. Examples

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include carbohydrate such as sucrose, lactose, mannitol, and dextran, proteins such as albumin and protamine, amino acids such as arginine, glycine, and threonine, surfactants such as Tween™ and Pluronic<sup>™</sup>, salts such as calcium chloride and sodium phosphate, and lipids such as fatty acids, phospholipids, and bile salts. The stabilizers are generally added to the protein in a ratio of 1:10 to 4:1, carbohydrate to protein, amino acids to protein, protein stabilizer to protein, and salts to protein; 1:1000 to 1:20, surfactant to protein; and 1:20 to 4:1, lipids to protein. Other stabilizers include high concentrations of ammonium sulfate, sodium acetate or sodium sulfate, based on comparative studies with heparinase activity. stabilizing agents, preferably the ammonium sulfate or other similar salt, are added to the enzyme in a ratio of 0.1 to 4.0 mg ammonium sulfate/IU enzyme.

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The use of stabilizers is demonstrated as follows. The purified chondroitinase enzymes were dialyzed into 10 mM sodium phosphate, pH 7.5, to a concentration of 2 IU/ml and supplemented with either 1 mg/ml bovine serum albumin, 1.5 M sodium acetate, 0.0025 M Tris or 0.15 M Tris, and an accelerated shelf life performed at 37°C. 2 IU of purified chondroitinase enzymes also were placed into various buffers, lyophilized and an accelerated shelf life performed at 37°C. The results are shown in Table 3.

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Table 3: Stability of chondroitinase enzymes at 37°C.
7 day retention of activity (%)

additive to	rmat chonc	iroiti	nase AC cnond	roitinase B
0.15 M Tris	liquid	1	42	
0.0025 M Tri	s liquid	22	44	
1 mg/ml BSA	liquid	1	26	
1.5 M NaOAc	liquid	64	72	
0.15 M Tris	lyophiliz	ed	26.7	43.7
PBS	lyophiliz	ed	8.7	15.9
8 mg/ml				
sucrose	lyophiliz	ed	88	93.16
2 mg/ml				
glycine	lyophiliz	ed	42.4	75.7

# Cloning of Chondroitinase AC and Chondroitinase B Amino Acid Analysis

The purified proteins were analyzed by the technique of Edman, Ann. N. Y. Acad. Sci. 88:602, 1950, to determine the N-terminal amino acid. 5 However, the Edman chemistry was unable to liberate an amino acid, indicating that a post-translational modification had occurred at the N-terminal amino acid of both chondroitinase proteins. One nmol 10 samples of chondroitinases AC and B were used for deblocking with pyroglutamate aminopeptidase. Control samples were produced by mock deblocking 1 nmol samples without adding the peptidase. All samples were placed in 10 mM ammonium carbonate 15 buffer at pH 7.5 with 10 mM dithiothreitol. 1 mU peptidase was added to the samples and the reaction allowed to incubate at 37°C for 8 hours. An additional 0.5 mU peptidase was added and incubation continued for 16 h. The reaction mixture was exchanged into 35 % formic acid by 20 diafiltration with 10,000 Dalton cut-off ultrafiltration membranes (Centricon, Amicon) and the sample dried under vacuum. Deblocked chondroitinase enzymes were then analyzed by Edman 25 chemistry to determine the N-terminal sequence, using an Applied Biosystems 745A Protein Sequencer.

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The N-terminal sequence of chondroitinase AC was QTGTAEL (Sequence ID No. 2, amino acids 24 to 30) and of chondroitinase B was VVASNEL (Sequence ID No. 4, amino acids 27 to 34).

The chondroitinase enzymes were subjected to enzymatic fragmentation using the arginine specific protease clostripain (EC 3.4.22.8, Sigma). Pre-activated clostripain was added to chondroitinase AC at a 1 to 2 % w/w ratio in 0.025 M sodium phosphate, 0.0002 M calcium acetate and 0.0025 M dithiothreitol at pH 7.5 ± 0.1 and incubated for 2 to 3 hours at 37°C. The reaction mixture was applied to a Vydac C<sub>18</sub> reverse phase HPLC column (0.46 cm I.D. x 30 cm) and the peptide fragments eluted at a linear flow rate of 1 cm-min<sup>-1</sup> with a linear gradient of 10 to 90 % acetonitrile in 1 % trifluoroacetic acid. Four of the peptide fragments obtained were subjected to amino acid sequence determination.

Clostripain was added to chondroitinase B at a 1 to 2 % w/w ratio in 0.025 M sodium phosphate, 0.0002 M calcium acetate and 0.0025 M dithiothreitol at pH 7.5 ± 0.1 and incubated for 2 to 3 hours at 37°C. The reaction mixture was applied to a Vydac<sup>TM</sup> C<sub>18</sub> reverse phase HPLC column and the peptide fragments eluted at a linear flow rate of 6.0 cmomin<sup>-1</sup> with a linear gradient of 10 to 90 % acetonitrile in 1 % trifluoroacetic acid. Three of the peptide fragments obtained were subjected to amino acid sequence determination.

## Construction of Flavobacterium heparinum gene library

A Flavobacterium heparinum chromosomal DNA library was constructed in lambda phage DASHII. 0.4  $\mu$ g of F. heparinum chromosomal DNA was partially digested with restriction enzyme, Sau3A, to produce a majority of fragments around 20 kb in size, as described in Maniatis, et al., Molecular Cloning, A

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laboratory Manual, 1982. This DNA was phenol/chloroform extracted, ethanol precipitated, ligated with DASHII arms and packaged with packaging extracts from a Lambda DASHII™/BamHI Cloning Kit (Stratagene, La Jolla, CA). The library was titered at approximately 10<sup>-5</sup> pfu/ml after packaging, was amplified to 10<sup>-8</sup> pfu/ml by the plate lysis method, and stored at -70°C as described by Silhavy et al. in Experiments with Gene Fusions, Cold Spring Harbor Laboratory, 1972.

The F. heparinum chromosomal library was titered to about 300 pfu/plate, overlaid on a lawn of E. coli, and allowed to transfect the cells overnight at 37°C, forming plaques. The phage plaques were transferred to nitrocellulose paper, and the phage DNA bound to the filters, as described in Maniatis, et al., ibid.

## Nucleic acid sequence encoding Chondroitinase AC

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Degenerate primers were designed from 20 peptides AC-1, AC-3 and AC-4 (Sequence ID No. 2, amino acids 395 to 413; 603 to 617; 514 to 536; and 280 to 288, respectively). Amplification of the primers was carried out in a 0.1 ml reaction buffer containing 50 mM KCl, 10 mM Tris/HCl pH 9, 0.1% 25 Triton X-100, 2.5 mM MgCl<sub>2</sub>, plus the four dNTPs at 200 μM, 2.5 units Tag Polymerase (Bio/Can, Mississauga, Ont.), 0.1 mM of each primer and 10 ng of F. heparinum genomic DNA. The amplified primers were linearized with SalI, NotI, and XbaI in 30 individual restriction digests, and combined, after purification, for use as template DNA. The samples were placed in an automated heating block, (DNA Thermocycler<sup>™</sup>, Barnstead/Thermolyne, Dubuque, IA) 35 programmed for cycles with temperatures of denaturation at 94°C for 1 min., annealing at 50°C for 2 min., and extension at 72°C for 2 min., with 35 repetitions of this sequence. The combination of

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synthetic oligonucleotide primers: 5'TCNGGRAARTARTANCCDATNGCRTCRTG-3' (Sequence ID No.
5), corresponding to peptide AC-3; and 5'TAYATGGAYTTYAAYGTNGARGG-3' (Sequence ID No. 6),
corresponding to peptide AC-4; yielded a PCR
product of approximately 750 bp in size. Attempts
to clone this fragment into vectors, pTZ/PC or into
pCRII (TA cloning kit, Invitrogen, San Diego, Ca.)
in E. coli strain, FTB1, were unsuccessful.

E. coli FTB1 was constructed as follows:
the F' episome from E. coli XL-1 Blue, (Stratagene,
La Jolla CA) carrying the lac Iq repressor gene was
moved, as described by Miller, Experiments in
Molecular Genetics, Cold Spring Harbor, 1972, into
E. coli TB1 described by Baker et al., Proc. Natl.
Acad. Sci. 81:6779-6783, 1984. The FTB1 background
permits a more stringent repression of
transcription from plasmids carrying promoters with
a lac operator such as the lac and tac promoters.

To facilitate cloning of these PCR products, a restriction site was incorporated at the 5' ends of the primers. The PCR products were analyzed for the absence of restriction sites which are found in the multiple cloning site of pBluescript (Stratagene, La Jolla, CA) to determine which restriction site should be added to the primers. This ensured that the PCR products would not be cut into multiple fragments when treated with the restriction enzyme used to form overhangs on the ends of the DNA fragments. BamHI met this criteria for all three PCR fragments. New primers were synthesized with BamHI sites at their 5' ends, which were otherwise identical to those described above, and used to produce a 764 bp PCR product, This DNA fragment was digested with Figure 1. BamHI, isolated on an agarose gel, as described by Maniatis et al., ibid, and purified using the

Geneclean™ kit (Bio/Can, Mississauga, Ont.) pBluescript was digested with BamHI, the 5' ends dephosphorylated by alkaline phosphatase treatment as described by Maniatis et al., ibid, and purified from an agarose gel using the Geneclean™ kit. 5 treated PCR fragment and pBluescript plasmid DNA were ligated, transformed into FTB1, and plated onto LB agar plates containing ampicillin at 0.2 mg/ml. Plasmids from colonies grown on these plates were isolated by colony cracking as 10 described in Maniatis et al., ibid. All enzymes were supplied by New England Biolabs (Mississauga, Ont.). Plasmids were isolated using the RPM™ kit (Bio/Can, Mississuaga, Ont.). Sequence analysis of the cloned PCR fragment correlated with reverse 15 transcribed peptide sequences from chondroitinase AC peptides, indicating that the PCR fragment encodes the chondroitinase AC gene. DNA sequencing was performed by the dideoxy-chain termination method of Sanger et al., Proc. Natl. Acad. Sci., 20 74:5463-5467, 1978. Sequencing reactions were carried out with the Sequenase™ Kit (U.S. Biochemical Corp., Cleveland, Ohio) and S-dATP (Amersham Canada Ltd., Oakville, Ontario, Canada), as specified by the supplier. 25

The 764 bp PCR fragment, contained in plasmid pA2C1BS-11 represents approximately 36% of the coding region for the Chondroitinase AC gene. This entire 764 bp fragment was sequenced and was found to contain a continuous open reading frame which encoded peptides AC-3, AC-4 and AC-1 (Sequence ID No. 2, amino acids 395-413; 603-617; 514-536; 280-288, respectively).

The 764 bp PCR fragment was used to probe
the genomic F. heparinase lambda library. First,
pA2C1BS-11 was isolated via the boiling method, as
described in Maniatis et al., ibid. The plasmid

was digested with BamHI, separated from the vector, purified as described above and labeled with a Nick Translation™ kit (Boehringer Mannheim, Montreal, Canada) using radiolabelled  $^{32}P$   $\alpha$ -dATP. E. coli P2392 (Stratagene, La Jolla, CA) was used as the 5 lawn for plating the lambda library. Approximately 6000 plaques were screened by plaque hybridization using BA85 nitrocellulose membranes (Scheicher & Schuell, Keene, NH) as described by Maniatis et al., ibid. Plague hybridization was carried out, at 10 65°C for 16 hours in a Tek Star™ hybridization oven (Bio/CAN Scientific, Mississauga, Ontario). Subsequent washes were performed at 65°C, twice for 15 min. in 2X SSC, once in 2X SSC/0.1% SDS for 30 min. and once in 0.5% SSC/0.1% SDS for 15 min. 15 More than 100 positive plaques were identified and isolated, some of which were clusters of plaques. These were rescreened by spotting the lambda clone onto a lawn of P2392 host cells and reprobing via 20 plaque hybridization. Six plaques were positive upon rescreening, and their DNA was isolated, as described by Maniatis, et al., ibid, and digested with restriction enzymes corresponding to the sites on the ends of lambda DASH II arms. This DNA was used in Southern hybridization analysis (Southern, 25 J. Mol. Biol. 98:503-517, 1975) by blotting onto Hybond™ N nylon membrane (Amersham, Oakville, Canada) using hybridization and wash conditions, described above for plaque hybridization. clone contained a 4.5 kb Sal I fragment and another 30 contained a 6 kb BamHI fragment, both of which hybridized with the probe. These were cloned into corresponding sites of pBluescript.

Because the molecular weight of

35 chondroitinase AC is approximately 75 kD, the size
of the corresponding gene would be approximately

2.05 kb. Both the 4.5 kb SalI and the 6 kb BamHI

chromosomal DNA fragments could include the entire chondroitinase AC gene. To increase the probability of analyzing a DNA fragment which encodes the entire gene, the 6 kb BamHI fragment was chosen for sequence analysis. The pBluescript plasmid containing this BamHI fragment (called p64BS2-7, Figure 1) was isolated using the Qiagene kit (Bio/Can, Miss, Ont). A method of DNA sequencing, the walking primer strategy (Voss et al. Meth. Molec. Cell. Biol. 3:153-155 (1992)), was 10 employed using synthetic primers (Eppendorf, model ECOSYN™ D300, Madison, WI) and an A.L.F. DNA sequencer (Pharmacia LKB, Mtl, Qc). Fluorescenated Universal and Reverse primers provided in the Pharmacia AutoRead kit were also used. 15 Fluorescently labeled dNTPs were incorporated into sequencing reactions with the Pharmacia AutoRead Fluorescent labelling kit (Pharmacia LKB, Mtl, QC). Areas of secondary structure were resolved by one of two methods. First, fluorescenated primers 20 which hybridized close to, and 5' to, the region of secondary structure were synthesized. Using these primers, the Pharmacia AutoCycle™ kit (Pharmacia LKB, Mtl, Qc), and a automated heating block (DNA Thermocycler™, Barnstead/Thermolyne, Dubuque, 25 Iowa), programmed for step cycles of 95°C for 36 sec, 50°C for 36 sec and 72°C for 84 sec, repeated 25 times, sequencing of secondary structure regions was accomplished. Any ambiguous areas still not resolved by the first method were sequenced by the 30 method of Sanger et al., Proc. Natl. Acad. Sci. 74: 5463-5467 (1978), using  $^{35}S$  a-dATP, and a USB Sequenase™ kit (LaJolla, Ca.) in which dGTP was replaced by dITP.

Analysis of the DNA sequence indicated that there was a single, continuous open reading frame of 2100 bp containing codons for 700 amino acid

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residues. All four clostropain-derived peptides were encoded by this gene. Searching for a possible signal peptide sequence using Geneworks™ (Intelligenetics, Mountain View, Ca.), suggested that there are two possible sites for the processing of the protein into a mature form: Q-23 (glutamine) and A-28 (alanine). N-terminal amino acid sequencing of deblocked, processed Chondroitinase AC indicated that the mature protein begins with Q-23 and contains 678 amino acids with a calculated molecular weight of 77,169 Daltons.

Expression of Chondroitinase AC in E. coli

Construction of an expression vector for chondroitinase AC is shown in Figure 2. The vector pGB is an *E. coli* expression vector which contains an unique *BamHI* site, whereby expression of a DNA fragment inserted into this site is driven by a double tac promoter. The vector also includes a kanamycin resistance gene and the *lac* I<sup>q</sup> gene to allow induction of transcription with IPTG. PCR was used to generate a mature chondroitinase AC gene.

An oligonucleotide, 5'GCGGATCCATGCAGCAGACCGGTACTGCAGAA-3', (Sequence ID
No. 7) was designed to insert an ATG-start site
immediately preceding the codon for the first amino
acid (Q-23) of mature chondroitinase AC, while an
oligonucleotide 5'-CGCGGATCCCCTAGATTACTACCATCAAAA3' (Sequence ID No. 8) was designed to hybridize
downstream of the TAG-stop codon. Both
oligonucleotides also contain a BamHI site.
Plasmid p64BS2-7 was used as the template in a PCR
reaction with an annealing temperature of 45°C. A
specific fragment of the expected size of 2034 bp
was obtained. This fragment was isolated and
inserted into a BamHI site of the expression vector
pGB.

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The construct was transformed into E. coli strain, F-TB1, and the transformed bacteria was grown at 37°C in LB medium containing 75  $\mu$ g/ml kanamycin to an  $OD_{600}$  of 0.5, at which point the tac promoter from pGB was induced by the addition of 1 mM IPTG. Cultures were grown an additional 2 to 5.5 hours at either 23°C, 30°C or 37°C. were cooled on ice, concentrated by centrifugation and resuspended in cold PBS at 1/10th the original culture volume. Cells were lysed by sonication and cell debris removed by centrifugation at 10,000 x g, 5 minutes. The pellet and supernatant fractions were analyzed separately for chondroitin sulfate A or C degrading (chondroitinase AC) activity. Chondroitin sulfate A degrading activities of 1.24  $\times 10^{-2}$ , 2.88  $\times 10^{-2}$ , and 4.25  $\times 10^{-2}$  IU/ml/OD and chondroitin sulfate C degrading activities of 1.57  $\times$  10<sup>-2</sup>, 2.24  $\times$  10<sup>-2</sup>, and 6.02  $\times$  10<sup>-2</sup> IU/ml/OD were observed from cultures grown at 23, 30 and 37°C, respectively. The activities using chondroitin sulfate A as the substrate are approximately twice that of those using chondroitin sulfate C as the substrate. This ratio is also observed when measuring the activity of the wild type chondroitinase AC using both these substrates.

E. coli F-TB1 (pGB-ChAC) was grown in a 3.5 L Braun Biostat E computer controlled fermenter in M9 medium to a dry cell weight concentration of 35 g/L. Glucose and ammonia were added as needed to maintain growth and pH at 7.0. Chondroitinase A activity accumulated to 103.44 IU/ml while chondroitinase C activity accumulated to 28.26 IU/ml.

## Nucleic Acid encoding Chondroitinase B

Partial-guessmer PCR primers were designed using the amino acid sequences of the clostripaingenerated peptides from the chondroitinase B

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> protein and the codons commonly found in Flavobacterium genes, Table 4. Three peptides were generated, designated CHB-1 (Sequence ID No. 4, amino acids 373 to 384), CHB-2 (Sequence ID No. 4,

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amino acids 41 to 50), and CHB-3 (Sequence ID No. 4, amino acids 130 to 146).

Table 4: Codon usage table for Flavobacterium and Escherichia coli. consensus codon

am	ino a	cid	codon	(s)	E. coli	Flavobacterium
			_			
A	GCT,	GCC,	GCG,	GCA	GCT	GCC
C	TGT,	TGC			EITHER	EITHER
D	GAT,	GAC			EITHER	EITHER
E	GAG,	GAA			GAA	GAA
	TTC,				EITHER	
G	GGC,	GGA,	GGG,	GGT	GGC or GGT	GGC
H	CAC,	CAT			CAT	CAT
I	ATC,	ATA,	ATT		ATA	ATC
K	AAA,	AAG			AAA	AAA
L	CTT,	CTA,	CTG,	TTG,		
	TTA,	CTC			CTG	CTG
M	ATG				ATG	ATG
N	AAC,	AAT			AAC	AAT
P	CCC,	CCT,	CCA,	CCG	CCG	CCG
	CAG,				CAG	CAG
R	CGT,	AGA,	CGC,	CGA,		
	AGG,				CGT	CGC
S	,		TCG,	TCT,		
	AGC,				TCT	ND
				ACA		ACC or ACA
		GTA,	GTT,	GTG		ND
W	TGG				TGG	TGG
Y	TAC,	TAT			EITHER	TAT

5'-CGG GAT CCC ARA TYG CCG AYG GNA CNT ATA AAG A-3' (Sequence ID No. 9) was derived from the CHB-2 peptide (Sequence ID No. 4, amino acids 41 to 50) and 5'-CGG GAT CCG GCN SKA TTG CGT TCR TCA AA-3' (Sequence ID No. 10) was derived from peptide CHB-3, Sequence ID No. 4, amino acids 130 to 146. A BamHI site was present on the 5' end of each primer to increase the efficiency of cloning of the PCR product. Using linear F. heparinum chromosomal DNA, described above, as a template, a single 300 bp DNA fragment was amplified. Conditions for the amplification were as follows:

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denaturation at 94°C for 40 sec, annealing at 45 or 50°C for 1 min. and extension at 72°C for 2 min. This cycle was repeated 35 times.

As shown in Figure 3, the PCR fragment was purified on an agarose gel, digested with BamHI and ligated into BamHI digested, dephosphorylated pBluescript. The ligation mixture was used to transform E. coli FTB1. Of the 50 resulting transformants, one yielded a 300 bp fragment when cut with BamHI. The insert in this plasmid, pCHB300, was subjected to DNA sequence analysis, performed as described above, which revealed that the insert contained DNA sequences outside of the primer regions which encoded amino acid sequence matching that determined for two chondroitinase B peptides. This insert was used to screen the lambda library of F. heparinum chromosomal DNA, which was constructed as described above.

The lambda library was plated with a density of 200 plaques per dish. Plate lifts of 20 dishes were made. For production of the probe, 500 ng of pCHB300 was submitted to 30 cycles of PCR amplification; denaturation at 93°C, annealing at 55°C and extension at 72°C, each for 1 min., using the primers described above. The resulting PCR fragment was purified on agarose gels and labelled with dATPa32P, using the Random Primer labelling kit (Boehringer Mannheim, Laval, Canada). potential lambda clones were found which hybridized with this probe, after the lifts were subjected to washing one time, in 2X SSC at 58°C. Rescreening of these plaques gave a positive signal for 17 of the plaques after washing at 58°C, 2X for 15 min. in 2X SSC, 1X for 30 min. in 2X SSC/0.1% SDS and 1X for 20 min. in 0.5X SSC/0.1% SDS. Two of 8 clones analyzed further showed a 5.0 kb HindIII fragment hybridizing with the probe and comigrating with a

HindIII fragment from F. heparinum chromosomal DNA which also hybridized with the 300 bp probe. 5.0 kb fragment was gel purified from both lambda clones, ligated into the HindIII site of pBluescript and transformed into FTB1.

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44 colonies were picked and rubbed on the side of a 0.5 ml PCR tube containing 20  $\mu$ l of the same PCR mixture as above. PCR was performed at: denaturation at 93°C, for 30 sec., annealing at 58°C, for 30 sec. and extension at 72°C, for 1 min, for 35 cycles. Upon analysis, 6 transformants showed amplification of the 300 bp band. DNA from these colonies were isolated and digested by HindIII revealing the presence of a 5.0 kb fragment. 5 out of the 6 clones hybridized with 15 the 300 bp fragment, confirming results of the PCR amplification experiment. One of these clones, pCHB78, was selected and used as a template for DNA sequencing.

Using a walking primer strategy, sequencing reactions were carried out as described above for the A.L.F. DNA sequencer. Sequence analysis revealed a single 1.52 kb open reading frame coding for 506 amino acid residues. The preprotein was found to have a signal peptide of 25 amino acids. The mature chondroitinase B enzyme contains 481 amino acids with a calculated molecular weight of 53,563 daltons.

## Expression of Chondroitinase B in E. coli

Construction of an expression vector for chondroitinase B is shown in Figure 4. Primers were designed to amplify the coding region of the chondroitinase B gene in an analogous manner to that described above with reference to expression of the chondroitinase AC gene. One oligonucleotide used for amplification of the chondroitinase B coding sequence (5'-

CGCGGATCCATGCAGGTGTTGCTCAAATGAAACT-3') (Sequence ID No. 11), contained a BamHI restriction site at its 5' end and an ATG codon that was to be inserted before the first amino acid of the mature protein. The second oligonucleotide (5'-CGGAATCAATTCACCGGG-AT-3') (Sequence ID No. 12) was designed with a XmmI restriction site and a termination codon to be inserted at the end of the coding sequence of the gene. Using 100 ng of pCHB78 as template, with an annealing temperature of 52°C, the 1.5 kb fragment was amplified, gel purified, restriction digested and inserted into pGB previously cut with BamHI and XmmI. This resulted in the definitive pGB-CHB construct used to express the protein.

This construct was transformed in E. colistrain DH5 $\alpha$ , expressed as described for the chondroitinase AC enzyme. After growing cells until an O.D. 600 = 0.5, 1 mM IPTG was added to the cultures to induce the tandem tac promoters and cells were transferred to either 23°C, 30°C or 37°C for additional growth for 5, 3 and 2 hours, respectively. After sonication, supernatant fractions were assayed for activity on dermatan sulfate. Growth of cells at 23°C gave the best results with a degrading activity of 0.57 IU/ml/OD while growth of cells at 30°C and 37°C gave degrading activities of 0.14 and 0.01 IU/ml/OD respectively.

The present invention describes a methodology for obtaining highly purified chondroitin degrading enzymes derived from the natural organism *Flavobacterium heparinum*, and the genes encoding these enzymes. Derivatives of the genes can be prepared by making conservative substitutions, additions and deletions thereof, which do not substantially impact on the resulting enzymatic activity, or by using degenerative forms

of the genes. As used herein, conservative substitutions involve substitutions of codons which encode the same amino acids and substitutions of amino acids for amino acids having similar structure or chemical characteristics, which are well known to those skilled in the art, for example, groups of structurally similar amino acids include (I,L,V); (F,Y); (K,R); (Q,N); (D,E); AND (G,A).

Variations of these methods will be obvious to those skilled in the art from the foregoing detailed description of the invention. Such modifications are intended to come within the scope of the appended claims.

# SEQUENCE LISTING

GENERAL INFORMATION: (1)

INC. APPLICANT: IBEX TECHNOLOGIES R AND D,

TITLE OF INVENTION: CHONDROITIN LYASE ENZYMES

NUMBER OF SEQUENCES: 19

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ZIP: 30309-3450 F

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

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B

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OPERATING SYSTEM: PC-DOS/MS-DOS

Version #1.25 PatentIn Release #1.0, SOFTWARE: ΰ 9

ATTORNEY/AGENT INFORMATION: B (viii)

REGISTRATION NUMBER: 31,284 NAME: Pabst, Patrea L.

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REFERENCE/DOCKET NUMBER: IT103

TELECOMMUNICATION INFORMATION: ์

(ix)

TELEPHONE: (404) 873-8794 B

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SEQUENCE CHARACTERISTICS: INFORMATION FOR SEQ ID NO:1:

Œ,

(2)

LENGTH: 2103 base pairs TYPE: nucleic acid  $\mathbf{g}$ (B)

STRANDEDNESS: single

TOPOLOGY: linear a

MOLECULE TYPE: DNA (genomic)

HYPOTHETICAL: NO

ANTI-SENSE: NO (i.v.)

FEATURE: (ix)

NAME/KEY: misc feature **A B** 

LOCATION:

(D) OTHER INFORMATION: /note= "Nucleic acid sequence encoding chondroitinase AC from Flavobacterium heparinum."
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

			!			
ATGAAGAAA	ATGAAGAAAT TATTTGTAAC CTGTATAGTC TTTTTCTCTA TTTTAAGTCC TGCTCTGCTT	CTGTATAGTC	TTTTCTCTA	TTTTAAGTCC	TGCTCTGCTT	9
ATTGCACAGO	ATTGCACAGC AGACCGGTAC TGCAGAACTG ATTATGAAGC GGGTGATGCT GGACCTTAAA	TGCAGAACTG	ATTATGAAGC	GGGTGATGCT	GGACCTTAAA	120
AAGCCTTTGC	AAGCCTTTGC GCAATATGGA TAAGGTGGCG GAAAAGAACC TGAATACGCT GCAGCCTGAC	TAAGGTGGCG	GAAAAGAACC	TGAATACGCT	GCAGCCTGAC	180
GGTAGCTGG	GGTAGCTGGA AGGATGTGCC TTATAAAGAT GATGCCATGA CCAATTGGTT GCCAAACAAC	TTATAAAGAT	GATGCCATGA	CCAATTGGTT	GCCAAACAAC	240
CACCTGCTAC	CACCTGCTAC AATTGGAAAC TATTATACAG GCTTATATTG AAAAAGATAG TCACTATTAT	TATTATACAG	GCTTATATTG	AAAAAGATAG	TCACTATTAT	300
GGCGACGAT	GGCGACGATA AAGTGTTTGA CCAGATTTCC AAAGCTTTTA AGTATTGGTA TGACAGCGAC	CCAGATTTCC	AAAGCTTTTA	AGTATTGGTA	TGACAGCGAC	360
CCGAAAAGCC	CCGAAAAGCC GCAACTGGTG GCACAATGAA ATTGCCACTC CGCAGGCCCT TGGTGAAATG	GCACAATGAA	ATTGCCACTC	CGCAGGCCCT	TGGTGAAATG	420
CTGATCCTG	CTGATCCTGA TGCGTTACGG TAAAAGCCG CTTGATGAAG CATTGGTGCA TAAATTGACC	TAAAAAGCCG	CTTGATGAAG	CATTGGTGCA	TAAATTGACC	480
GAAAGAATG	GAAAGAATGA AGCGGGGCGA ACCGGAGAAG AAAACGGGGG CCAACAAAAC AGATATCGCC	ACCGGAGAAG	AAAACGGGGG	CCAACAAAAC	AGATATCGCC	540
CTGCATTACT	CTGCATTACT TITAICGIGC TITGITAACG ICTGAIGAGG CITIGCIIIC CITCGCCGIA	TTTGTTAACG	TCTGATGAGG	CTTTGCTTTC	CTTCGCCGTA	009
AAAGAATTGI	AAAGAATTGT TTTATCCCGT ACAGTTTGTA CACTATGAGG AAGGCCTGCA ATACGATTAT	ACAGTTTGTA	CACTATGAGG	AAGGCCTGCA	ATACGATTAT	099
TCCTACCTG	TCCTACCTGC AGCACGGTCC GCAATTACAG ATATCGAGCT ACGGTGCCGT ATTTATTACC	GCAATTACAG	ATATCGAGCT	ACGGTGCCGT	ATTTATTACC	720
GGGGTACTG	GGGGTACTGA AACTTGCCAA TTACGTTAGG GATACCCCTT ATGCTTTAAG TACCGAGAAA	TTACGTTAGG	GATACCCCTT	ATGCTTTAAG	TACCGAGAAA	780
CTGGCTATA	CTGGCTATAT TTTCAAAGTA TTACCGCGAC AGTTATCTGA AAGCTATCCG TGGAAGTTAT	TTACCGCGAC	AGTTATCTGA	AAGCTATCCG	TGGAAGTTAT	840
ATGGATTTT	ATGGATTTTA ACGTAGAAGG CCGCGGAGTA AGCCGGCCAG ACATTCTAAA TAAAAAGGCA	CCGCGGAGTA	AGCCGGCCAG	ACATTCTAAA	TAAAAAGGCA	900
GAAAAAAG	GAAAAAAGA GGTTGCTGGT GGCGAAGATG ATCGATCTTA AGCATACTGA AGAATGGGCT	GGCGAAGATG	ATCGATCTTA	AGCATACTGA	AGAATGGGCT	960

2040	AGGAAAAGA	ATTTAAAAAC	じじせるしたない	אטאפאסאסר אפראפייייים ארייבייייים ארייביייסר ארייבייקרארייבייי אניביבעבעבע מבמיבייייניי	טפטפפפטפפ מפטפפ	女女 ないじむしないし
1980	GGCTGCCGAT	AGGTAATTTG	AATGGCAAGC	AAGCCATGTG CAGTGCTGAT CAAGCACATC AATGGCAAGC AGGTAATTTG GGCTGCCGAT	CAGTGCTGAT	AAGCCATGTG
1920	TGAAACAGAT	GCATAGAAAT	AGCGTAGCGG	CAGGCTATCT TCTATACAGC TGGAAAATTA AGCGTAGCGG GCATAGAAAT TGAAACAGAT	TCTATACAGC	CAGGCTATCT
1860	AGATATGGTA	ATCAGCAGTT	GCAGITIAIC	AAAGTCCTTG CCAATACCAA CCAGCTGCAG GCAGTTTATC ATCAGCAGTT AGATATGGTA	CCAATACCAA	AAAGTCCTTG
1800	AACGGCACCG	AATATAATGG	GAAATTAAAA	ATCGITITGC CGGGAATAAA CAAGCCGGAA GAAATTAAAA AATATAATGG AACGGCACCG	CGGGAATAAA	ATCGTTTTGC
1740	GTATGCTTAT	AAAATGCGCA	GCCAGGCCAG	GATGTATTTA AGCTTTGGAT CAACCATGGT GCCAGGCCAG	AGCTTTGGAT	GATGTATTTA
1680	AGTTTCTGGT	CAAAAGATGA	AATTCACATT	TCGCAAAAAG GCAATTGGTT CCACATCAAC AATTCACATT CAAAAGATGA AGTTTCTGGT	GCAATTGGTT	TCGCAAAAAG
1620	GAGTACCCAG	ACCTTAGTCT	GAAGGGGCCA	TIGCACGAIG CGAITGGITA TTACTITCCT GAAGGGGCCA ACCTTAGICT GAGIACCCAG	CGATTGGTTA	TTGCACGATG
1560	GTTCTGGTTG	CACAGGGACA	ACGTTTAAAG	GCAGGTAAAA CCGGCCGGGG TAAAATAACA ACGTTTAAAG CACAGGGACA GTTCTGGTTG	ອອອອລລອອລລ	GCAGGTAAAA
1500	TATAAGTACT	ATGGCCCGGT	AGCTGGTTAA	CCTGAAAACA TTACCACTAC CCTTAACCAG AGCTGGTTAA ATGGCCCGGT TATAAGTACT	TTACCACTAC	CCTGAAAACA
1440	CAGCAATGCC	CCGGTATCAA	TGTCTTGGTG	GCCTGGTTCT TTTTGACAA AGAGATTGTA TGTCTTGGTG CCGGTATCAA CAGCAATGCC	TTTTGACAA	GCCTGGTTCT
1380	GGCAAAGAAA	ATAGCTTACA	TTGGATTACG	GATGGTGTAT ACGGGGCCAG TGCCTACGCA TTGGATTACG ATAGCTTACA GGCAAAGAAA	ACGGGGCCAG	GATGGTGTAT
1320	AGGGGTGTCT	ACTTTGCAGG	GGGAGCAATG	AGACCITIGA CGAAGCITIG GGGAGAGCAG GGGAGCAAIG ACTITGCAGG AGGGGIGICI	CGAAGCTTTG	AGACCTTTGA
1260	TTTAACCGAC	GCCGTGATTA	GGCATAACCA	AIGCCGGIAT GGGAAIGGGA CAAGAIICCI GGCAIAACCA GCCGIGAIIA IITAACCGAC	GGGAATGGGA	ATGCCGGTAT
1200	CTATAACATT	GACCAGAATA	CAATTGCGCG	AGGTATITAT CTGATGGGGC TACTAACATA CAATTGCGCG GACCAGAATA CTATAACATT	CTGATGGGGC	AGGTATTTAT
1140	ccrecresec	ATAAAGAAAA	GAATCCGGCA	CGTATGGTGA GTAAGCGGAC CCGACGCAGT GAATCCGGCA ATAAAGAAAA CCTGCTGGGC	GTAAGCGGAC	CGTATGGTGA
1080	TTTTAATGTT	CTGCCTATTC	CATTTAAGAC	CATCAGITCI GGAAIGGIGA ITAIGIGCAA CAITIAAGAC CIGCCIAIIC IITIAAIGII	GGAATGGTGA	CATCAGTTCT
1020	GCCCTATCAC	ATAAGATTGA	GCGGCCGGCT	GATGCGATAG CCAGGACAGA TAGCACAGTT GCGGCCGGCT ATAAGATTGA GCCCTATCAC	CCAGGACAGA	GATGCGATAG

AATCGGGTAA AAATTGATTT TCCGCAACAG GAATTTGCAG GTGCAACGGT TGAACTGAAA 2100
TAG 2103
(2) INFORMATION FOR SEQ ID NO:2:     (i) SEQUENCE CHARACTERISTICS:     (A) LENGTH: 700 amino acids     (B) TYPE: amino acid     (D) TOPOLOGY: linear     (ii) MOLECULE TYPE: protein     (ix) FEATURE:     (A) NAME/KEY: Peptide     (B) LOCATION: 123     (D) OTHER INFORMATION: /note= "Amino acids 1 through 23 are a leader     peptide."     (ix) FEATURE:     (ix) REATURE:     (ix) NAME/KEY: misc_feature     (ix) REATURE:     (ix) OTHER INFORMATION: /note= "Amino acid sequence     (ix) PEATURE:     (ix) PEATURE:     (ix) NAME/KEY: misc_feature     (ix) REATURE:     (ix) PEATURE:     (ix) NAME/KEY: misc_feature     (ix) NAME/KEY: misc_feature     (ix) PEATURE:     (ix) NAME/KEY: misc_feature     (ix) PEATURE:     (ix) PEATURE:     (ix) REATURE:     (ix) PEATURE:     (ix) PEATURE:
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
Met Lys Lys Leu Phe Val Thr Cys Ile Val Phe Phe Ser Ile Leu Ser  1 15 16 25 17 18 26 19 19 10 15 10 26 10 27 10 28 29 10 29 20 20 20 20 20 20 20 20 20 20 20 20 20

Lys	Asn 80	Asp	Ala	His	Met	Thr160	Lys	Asp	Gln	Gln	Thr 240	Leu	Tyr	Arg	Arg	Ala 320
Trp	Asn	Lys 95	Lys	Trp	Leu	Leu	Asn 175	Ser	Val	Leu	Ile	Ala 255	Ser	Gly	Lув	Trp
Ser	Pro	Glu	Ser 110	Trp	Ile	Lys	Ala	Thr 190	Pro	Tyr	Phe	Tyr	Asp 270	Glu	Lys	Glu
Gly	Leu	Ile	Ile	Asn 125	Leu	His	$_{ m Gly}$	Leu	Tyr205	Ser	Val	Pro	Arg	Val 285	Glu	Glu
Asp 60	Trp	Tyr	Gln	Arg	Met 140	Val	Thr	Leu	Phe	<b>TYr</b> 220	Ala	Thr	Tyr	Asn	Ala 300	$\mathtt{Thr}$
Pro	Asn 75	Ala	Asp	Ser	Glu	Leu 155	Lys	Ala	Leu	Asp	Gly 235	Asp	Tyr	Phe	Lув	His 315
Gln	$\operatorname{Thr}$	Gln 90	Phe	Lys	Gly	Ala	Lys 170	Arg	Glú	Tyr	Tyr	Arg 250	Lys	Asp	Lys	Lys
Leu	Met	Ile	Val 105	Pro	Leu	Glu	Glu	Tyr 185	Lys	Gln	Ser	Val	Ser 265	Met	Asn	Leu
$\operatorname{Thr}$	Ala	Ile	Lys	Asp 120	Ala	Asp	Pro	Phe	Val 200	Leu	Ser	Tyr	Phe	<b>Tyr</b> 280	Leu	Asp
Asn 55	Asp	Thr	Asp	Ser	Gln 135	Leu	Glu	Tyr	Ala	G1y 215	Ile	Asn	Ile	Ser	11e 295	
Leu	Asp 70	Glu	Asp	Asp	Pro	Pro 150	$_{ m G1y}$	His	Phe	Glu	Gln 230	Ala	Ala	Gly	Asp	Met 310
Asn	Lys	Leu 85	$_{ m G1y}$	Tyr	Thr	Lys	Arg 165	Leu	Ser	Glu	Leu	Leu 245	Leu	Arg	Pro	Lys
Lys	Tyr	Gln	<b>Tyr</b> 100	Trp	Ala	Lys	Lув	Ala 180	Leu	Tyr	Gln	Lys	<u>Гув</u> 260	Ile	Arg	Ala
Glu	Pro	Leu	Tyr	$\frac{Tyr}{115}$	Ile	Gly	Met	Ile	Leu 195	His	Pro	Leu	Glu	Ala 275		Val
Ala 50	Val	Leu	His	Lys	Glu 130	Tyr	Arg	Asp	Ala	Val 210	Gly	Val	Thr	Lys	Val 290	• •
Val	Asp 65	His	Ser	Phe	Asn	Arg 145	Glu	Thr	Glu	Phe	His 225	Gly	Ser	Leu	Gly	Leu 305

Ala Ile Phe Arg TyrGly Lys Ser Asn Gln Tyr Gln 430 Ala Ser Asn Thr 510Val Glu Glu 590 Ser Gln Ser Ala  $_{\rm Ile}$ 11e 525 Glu Pro G1y 445 Pro G1yThr  $_{
m Glu}$ Val Lys Ala Gln 540 Lys 460 Asn Lys Ala Glu Ile TyrIle TyxG1ySer G1y 380 Lys 555 Ala Ser  $_{
m G1y}$ Asp Pro 395 Gly Lys G1y 475  $\operatorname{Thr}$ Val Len Val Ile G1yHis Met Leu Pro 410 Leu Ala Ala Gln 490 Arg Ser Ser  $_{
m G1y}$ G1y 570 Arg Leu His His Arg Lys 425 Asp Gln Leu Gly 585  $\operatorname{Thr}$ Asn 345 Asn  $_{\rm Ile}$ Gly Asn G1y 505 Ser Asn Len Ser 440 Len Lea Ser Trp Val 360 Glu Lys  $\operatorname{Thr}$ Thr Leu 520 Leu Pro Lys 375 Gln Asp Cys Lys Leu 535 Asn Len Thr Trp Phe Asn Val Ser 455 Trp  $\operatorname{Thr}$ Phe Trp Asn 11e 390 Pro GlyAsp GlyAsn Asn 550 Val 470 Leu 565 Ile Ala Ala  $^{
m Thr}_{485}$ Ile Glu 405 Arg G1yTyr Ile G1yAsn  $_{
m G1y}$ His Lys Asp Ile Gly Thr 500 Ala Ser Thr Trp His 340 Ala Leu Lys Asn Gln 515 Glu Ala Phe 435 Ile Ser Val Thr Tyr Pro 530  $\mathbf{T}\mathbf{yr}$ Ala 450 Ala Asp Gln Phe Tyr

Lys Lys Tyr Asn Gly Thr Ala Pro Lys Val Leu Ala Asn T 595		Thr	Asn G	Gln
Val Tyr His Gln Gln Leu Asp Met Val Gln 610		Ala	Ile P	Phe
Ala Gly Lys Leu Ser Val Ala Gly Ile Glu Ile 630		Glu	Thr A	Asp : 640
Pro Cys Ala Val Leu Ile Lys His Ile Asn Gly Lys 645		Gln	Val I 655	Ile
Leu Gln Lys Glu Lys Thr Ala Val 665		Leu 670	Ser I	Ile
Thr Gly Lys Thr Asn Arg Val Lys Ile 685		Asp	Phe P	Pro
Phe Ala Gly Ala 695	27.8 700			
<pre>(2) INFORMATION FOR SEQ ID NO:3:     (i) SEQUENCE CHARACTERISTICS:         (A) LENGTH: 1521 base pairs         (B) TYPE: nucleic acid         (C) STRANDEDNESS: single         (D) TOPOLOGY: linear         (ii) MOLECULE TYPE: DNA (genomic)         (iii) HYPOTHETICAL: NO         (iv) ANTI-SENSE: NO         (iv) ANTI-SENSE: NO         (ix) FEATURE:         (A) NAME/KEY: misc_feature         (B) LOCATION: 11521         (D) OTHER INFORMATION: /note= "Nucleotide sequence encoding chondroitinase B from flavobacterium</pre>	de sequ	ence teriu	E	
heparinum. " (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:				
ATGAAGATGC TGAATAAACT AGCCGGATAC TTATTGCCGA TCATGGTGCT GCTGAATGTG	GTGCT G	SCTGAR	TGTG	9
GCACCATGCT TAGGTCAGGT TGTTGCTTCA AATGAAACTT TATACCAGGT TGTAAAGGAG	CAGGT 1	rgtaar	GGAG	120
TCAGCTGATT TCAGCTGATT	GATGT T	rcagci	GATT	180

1320	AAAATTAAA	ATAAGCCGGT	AGGICIGCCI	AAGGGAATCC CTGTTAACAT TTCGGCCAAT AGGTCTGCCT ATAAGCCGGT AAAAATTAAA	CTGTTAACAT	AAGGGAATCC
1260	AGGTGTGGAA	TGGACTGGTA ATGTAGCCTT AGGTGTGGAA	TGGACTGGTA	GAAAAATAGC	TTATAGCAGG	GATGATTATT
1200	ATTTTTAAA	TTTCTTTAAG GATAAACCTT ATGTTTACCC ATTTTTTAAA	GATAAACCTT		CTAATGTTAA AAGGCAATCT	CTAATGTTAA
1140	CCCGCACCAG	TTGTGCAGCC AATAGGCTTA AGTTCGAAAC	AATAGGCTTA		GATGAGCGCA GAAAAGAATA	GATGAGCGCA
1080	TAATCCATTG	CCATCCATTT	AATGGGTATG	CATCAATGTA AATGGGTATG	TTGATAGCCA ACAACGCTTT	TTGATAGCCA
1020	TTTCGATATG	GCTTCGGAGC ATGCTCTTGC		ATTTAAACCC CGGTGCTATG	ATTTAAACCC	GCCGCATTGT
096	GAGGGGGAAC	CTGTCCGAAA CCATAAAGTC		TTATTTGAG	TAGCCTGTAA	AGGCATGTCA
006	TTGGGGAAGC	GGATACGGGG GAATGTTTGT	GGATACGGGG	CCAGCGATTT	TAGGCAATGA	AATTTTATA
840	GGCCATTAAC	CGTCACGGTG ATCATCAGGT		CATGAACTTT	GCCAGGGCAC	TACCTGAATT
780	TGGTAATACT	TCGCAGGAAA ATGTTTATTA	TCGCAGGAAA	CACCAGCAAA	CAGAGATCAT	GATTCGGAAG
720	TATGCGTCAG	CTAACCTGTT	CTGGTAGACT	AGGCCGTTGT	GTAATGATAT	GGCTATTACC
099	AATCAGGATT	CCGGAGGGGG	CCGGGTAATG	TTTTTTCCAA TCCGCAAAAA CCGGGTAATG	TTTTTCCAA	GATCACTGTT
009	CCATCGTGTT	CGGGGATGTA	GTGGGAGGAC	AATACAGCCA GAGCTATTAA AGACGGTTCG	GAGCTATTAA	AATACAGCCA
540	TAACCTGAAC	ACCAGGTAAT	ATCACTTTTG	TACCGATAAG ATCACTTTTG	ATTGCAGTTT	CGCATAGACC
480	TCAACATTGC	ACCGAAGACG GAAAGGTACC		TACTTCGCTT	CTTACATTAC	GCCAATTCTG
420	TTTTGATGAA	TATTTGATTG	ACCGCATGTG	TAACCGCATT ACCGCATGTG	ATGGTAGCTA	GTGGCTATAT
360	ACCCGGATTG	CAGGCATGGA AATCACATGG	CAGGCATGGA	CAGAGCTATT	AAGACGGGAA	ATCTGGTTTA
300	ACTGGAAGGC	CTGAGGGGG AGCACCTGAT	CTGAGGGGCG	TAAAGTAGAG	CCGGAGATGC	GTTTTTTTA
240	CCCGGGTAAG	AAGCCCTGAA	ATCACTATTA	GTCAGCAATT CAGGAAAATC TGGTTTGCCC ATCACTATTA AAGCCCTGAA CCCGGGTAAG	CAGGAAAATC	GTCAGCAATT

His  $_{\rm Gln}$ Ile Arg Tyr Gln Lys 80 Leu Ala Asn Ala Gly Pro Ser Val 255 Arg His 95 Ser His Gln 175 TyrArg His Val Val G1ySer Asn Gln Met Asn Glu 11e 110 Ser Asp Ser 190 Glu Arg Phe 205 Gly Ala Phe Asn G1y 285 Pro  $_{\rm Gly}$ Val Leu Asn Gly Gln Arg Ile Phe Met Arg Leu G1yIle 60 Glu Val Asp 11e 220 Tyr Ala 75 Leu LysCys Asn 235 Ser Tyr Lys 155 Arg Glu Val Lys Pro Gln Leu Asn Ile Trp Gly Phe Asp His  $_{\rm Ile}$ Glu 90 Gly Ala Lys 170 Ser Lys 250 LysG1yAla 185 Asp Gln 265 Asn Val Cys Asp Asp  $_{
m G1y}$ Val Ile Val Asp 105 Val Asp Ser Asp Asn 280 Phe Lys Glu Arg Val 200 Gly Thr Lys Leu 120 Asp  $\operatorname{Thr}$ Val  $\operatorname{Thr}$ Cys Leu Ile Ala Phe Ala Ile Ile Asn Ile Met 295 Lys Phe Phe 135 Thr Lys 55 G1yG1y 215 Tyr His Arg Asp  $\operatorname{Thr}$ Len Ala Trp Pro Leu 150 Ser Ala Сув 230 Tyr Pro 70 Val Gly Gly Gln Val Val Asn Asn Glu 245 TyrArg Gly 85 Ile Cys Cys 165 Len Gly  $\operatorname{Thr}$ Ser Asn 180 Met His G1yAla Thr 260 Gln His  $\operatorname{Thr}$ GlyGly 100 Asp Lea G1y 195 Glu Asn His 275 Tyr 35 Asp Ser 115 Thr Phe Glu Asp G1yAsp Phe 290 Asn Lys Ala 50 Gln Arg Gly Trp G1y 65 Val

Ile Cys Lys Lys 400 Ala Ser Pro G1y 480 Phe Gly Len Lys Leu Arg G1yPro G1yAla Asn TyrPhe Lys 495 Glu Val Glu Ala Arg 510 Glu Met Asn Gly Met His Phe Asn 430 Ser Arg Val 350 Glu Pro Ala Gly $_{\rm Gly}$ Glu Leu Ala Ile Thr Asn Lys 365 TyrLys Arg Ile Ile Lys Ser Ile Arg Gln Ťhr Ser Pro 380 460 Asp Phe Gln Ile Leu Ile Ala Phe Arg His Val 395 Trp Gly His Trp Ala Aen Ile Ala LysGlu Pro Ser  $\operatorname{Thr}$ Met  $\mathbf{I}$ 410 330 Asp Glu Gla Ser TyrAla Arg Leu Ser Asn 345 Ala Asp Thr Asp Lys Pro Lys Asn Val Pro Glu Lys  $_{\rm Ile}$ Leu Pro Glu Leu Ser Gly Lys Arg Asn Lys Ile Ala Asn 360 GlyArg Ile Len Asn Pro Phe Asn Pro Phe Lys Ile 455 Lys Lys Ala  $_{\rm G1y}$ Ala Glu Val Lys Val  $\operatorname{Thr}$ Leu Leu Phe Ile Phe Leu Leu Asn 405 Pro Asp Ile Thr Tyr Ala Leu Phe Glu His Arg Phe Met Tyr Tyr 420 Asp Trp Asn Leu Lys Asn Cys Asn Asp Ile  $\mathbf{T}\mathbf{y}\mathbf{r}$ Val Ala Leu Ala Val Gly Ser TyrLen Phe Ala Ala Asp Asp 450 G1y 385 Ala Ala Lea Lys Leu 465 Ala Ala Ala Ala

INFORMATION FOR SEQ ID NO:5: (3)

LENGTH: 29 base pairs SEQUENCE CHARACTERISTICS

TYPE: nucleic acid <u>(a)</u>

STRANDEDNESS: single

TOPOLOGY: linear 9

MOLECULE TYPE: DNA (synthetic)

HYPOTHETICAL: NO

ANTI-SENSE: NO (ix)

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- NAME/KEY: misc\_feature (B)
  - LOCATION: 1..29
- OTHER INFORMATION: /note= "Nucleotide sequence encoding peptide AC-3." SEQUENCE DESCRIPTION: SEQ ID NO:5: (xi)

# TCNGGRAART ARTANCCDAT NGCRICRIG

(2)

29

INFORMATION FOR SEQ ID NO:6:

- SEQUENCE CHARACTERISTICS:  $\widehat{\mathbb{A}}$ 
  - LENGTH: 23 base pairs (B)
    - TYPE: nucleic acid
- STRANDEDNESS: single <u>ပ</u>
- MOLECULE TYPE: DNA (synthetic) TOPOLOGY: linear <u>e</u>
  - HYPOTHETICAL: NO
    - ANTI-SENSE: NO <u>†</u>
      - FEATURE:
- NAME/KEY: misc feature LOCATION: 1..23 <u>8</u>
  - (B)
- OTHER INFORMATION: /note= "Nucleotide sequence encoding peptide AC-4." SEQUENCE DESCRIPTION: SEQ ID NO:6: 9 (xi)

# TAYATGGAYT TYAAYGTNGA RGG

(2) INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS:

- LENGTH: 32 base pairs (B)
  - TYPE: nucleic acid
- STRANDEDNESS: single Û
  - TOPOLOGY: linear
- MOLECULE TYPE: DNA (synthetic) 9
  - HYPOTHETICAL: NO
    - ANTI-SENSE: NO

      - FEATURE:
- (A) NAME/KEY: misc\_feature (B) LOCATION: 3.8

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ω (D) OTHER INFORMATION: /note= "Nucleotides 3 through encode a BamHI site."

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

GCGGATCCAT GCAGCAGACC GGTACTGCAG AA

INFORMATION FOR SEQ ID NO:8: (3)

SEQUENCE CHARACTERISTICS:

LENGTH: 30 base pairs TYPE: nucleic acid (B)

STRANDEDNESS: single

<u>U</u>

(D) TOPOLOGY: linear MOLECULE TYPE: DNA (synthetic)

(ii)

HYPOTHETICAL: NO (iii)

ANTI-SENSE: NO

FEATURE: (ix)

NAME/KEY: misc feature

LOCATION: 4..9

OTHER INFORMATION: /note= "Nucleotides 4 through (B) 9

SEQUENCE DESCRIPTION: SEQ ID NO:8: encode a BamHI site."

(xi)

CGCGGATCCC CTAGATTACT ACCATCAAAA

INFORMATION FOR SEQ ID NO:9: (i) SEQUENCE CHARACTERISTICS:

(2)

LENGTH: 34 base pairs (A)

TYPE: nucleic acid (B)

STRANDEDNESS: single

TOPOLOGY: linear <u></u> <u>a</u>

MOLECULE TYPE: DNA (synthetic)

HYPOTHETICAL: NO

ANTI-SENSE: NO (ix)

FEATURE:

NAME/KEY: misc feature **A** (B)

LOCATION:

 (A) NAME/KEY: misc feature
 (B) LOCATION: 1..29
 (D) OTHER INFORMATION: /note= "Nucleotide sequence derived from the CHB-3 (D) OTHER INFORMATION: /note= "Nucleotide sequence derived from the CHB-2 34 ω OTHER INFORMATION: /note= "Nucleotides 3 through peptide." encode a BamHI site." SEQUENCE DESCRIPTION: SEQ ID NO:9: MOLECULE TYPE: DNA (synthetic) CGGGATCCCA RATYGCCGAY GGNACNTATA AAGA NAME/KEY: misc\_feature (A) LENGTH: 29 base pairs STRANDEDNESS: single (2) INFORMATION FOR SEQ ID NO:10: (1) SEQUENCE CHARACTERISTICS: TYPE: nucleic acid TOPOLOGY: linear LOCATION: 3..8 HYPOTHETICAL: NO ANTI-SENSE: NO FEATURE: FEATURE: <u>0</u>0 (B) **(4)** <u>e</u> e (xi) (ix) (ix) (iv)

OTHER INFORMATION: /note= "Nucleotides 3 through 8 encode a BamHI (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: (A) NAME/KEY: misc\_feature
(B) LOCATION: 3..8 <u>@</u> site."

(2) INFORMATION FOR SEQ ID NO:11:

CGGGATCCGG CNSKATTGCG TTCRTCAAA

FEATURE:

peptide." (ix) I (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs

```
TYPE: nucleic acid
```

STRANDEDNESS: single Û

TOPOLOGY: linear 9

MOLECULE TYPE: DNA (synthetic)

HYPOTHETICAL: NO ANTI-SENSE: NO

FEATURE:

NAME/KEY: misc\_feature (F)

LOCATION: 1..34

OTHER INFORMATION: /note= "Oligonucleotide used for amplification of <u>e</u> (e)

the chondroitinase B coding sequence." (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

CGCGGATCCA TGCAGGTGTT GCTCAAATGA AACT

34

SEQUENCE CHARACTERISTICS: INFORMATION FOR SEQ ID NO:12:

(5)

LENGTH: 18 base pairs

TYPE: nucleic acid STRANDEDNESS: single

(C)

TOPOLOGY: linear

MOLECULE TYPE: DNA (synthetic) 9

HYPOTHETICAL: NO

ANTI-SENSE: NO

SEQUENCE DESCRIPTION: SEQ ID NO:12: (yt) (X1)

CGGAATCAAT TCACCGGG

We claim:

- 1. A purified chondroitinase degrading enzyme isolated from bacteria.
- 2. The enzyme of claim 1 selected from the group consisting of chondroitinase AC and chondroitinase B from Flavobacterium heparinum.
- 3. The enzyme of claim 2 expressed in bactera from a gene isolated from Flavobacterium heparinum.
- 4. The enzyme of claim 2 where the enzyme is chondroitinase AC and has a molecular weight between 72,000 and 82,000 Daltons and is capable of degrading chondroitin sulfate A and chondroitin sulfate C.
- 5. The enzyme of claim 2 where the enzyme is chondroitinase B and has a molecular weight between 52,700 and 57,300 Daltons and is capable of degrading dermatan sulfate or chondroitin sulfate B.
- 6. The enzyme of claim 4 encoded by the nucleotide sequence of Sequence ID No. 1 and sequences having conservative or degenerative substitutions thereof.
- 7. The enzyme of claim 4 having the amino acid sequence of Sequence ID No. 2 and sequences having conservative substitutions thereof.
- 8. The enzyme of claim 5 encoded by the nucleotide sequence of Sequence ID No. 3 and sequences having conservative or degenerative substitutions thereof.
- 9. The enzyme of claim 5 having the amino acid sequence of Sequence ID No. 4 and sequences having conservative substitutions thereof.
- 10. The enzyme of claim 1 further comprising a pharmaceutically acceptable carrier.

- 11. An isolated nucleotide sequence encoding an enzyme selected from the group consisting of chondroitinase AC and chondroitinase B from Flavobacterium heparinum.
- 12. The sequence of claim 11 naturally occurring in Flavobacterium heparinum.
- 13. The sequence of claim 11 where the enzyme is chondroitinase AC and has a molecular weight between 72,000 and 82,000 Daltons and is capable of degrading chondroitin sulfate A and chondroitin sulfate C.
- 14. The sequence of claim 11 where the enzyme is chondroitinase B and has a molecular weight between 52,700 and 57,300 Daltons and is capable of degrading dermatan sulfate or chondroitin sulfate B.
- 15. The sequence of claim 4 having the nucleotide sequence of Sequence ID No. 1 or sequences having conservative or degenerative substitutions thereof.
- 16. The sequence of claim 13 encoding the amino acid sequence of Sequence ID No. 2 or sequences having conservative substitutions thereof.
- 17. The sequence of claim 14 having the nucleotide sequence of Sequence ID No. 3 or sequences having conservative or degenerative substitutions thereof.
- 18. The sequence of claim 14 encoding the amino acid sequence of Sequence ID No. 4 or sequences having conservative substitutions thereof.
- 19. A method for purifying a chondroitin lyase from bacteria comprising:

lysing the bacteria;

extracting proteins from the periplasmic space of the lysed bacteria;

separating the extracted proteins by cation exchange chromatography using a salt or pH gradient;

separating the fractions having enzymatic activity obtained by elution of the cation exchange chromatography matrix by chromatography on a sulfated cellulose resin using a salt or pH gradient;

separating the fractions having enzymatic activity obtained by elution of the sulfated cellulose resin on hydroxyapatite using a salt or pH gradient;

separating the fractions having enzymatic activity obtained by elution of the hydroxyapatite by chromatography using cation exchange chromatography using a salt or pH gradient; and

separating the fractions with enzymatic activity on the basis of molecular weight.

FIG. 1

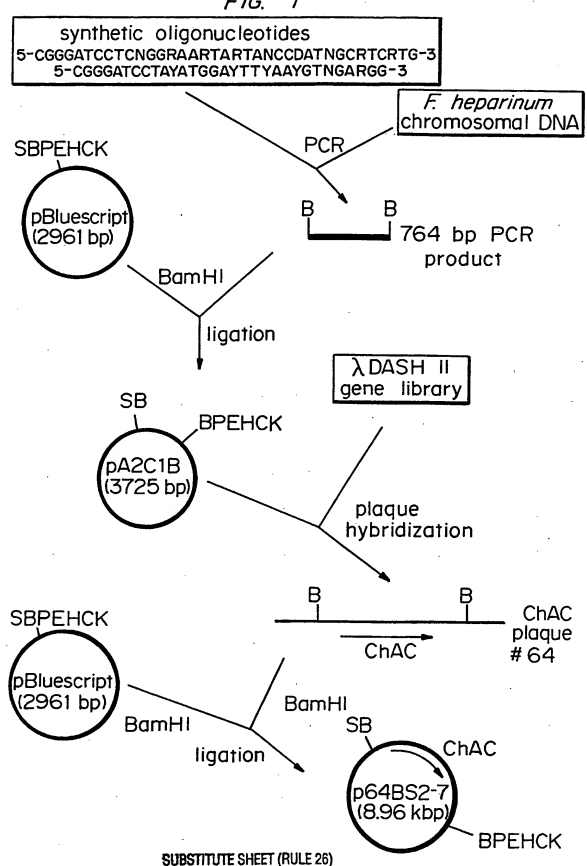
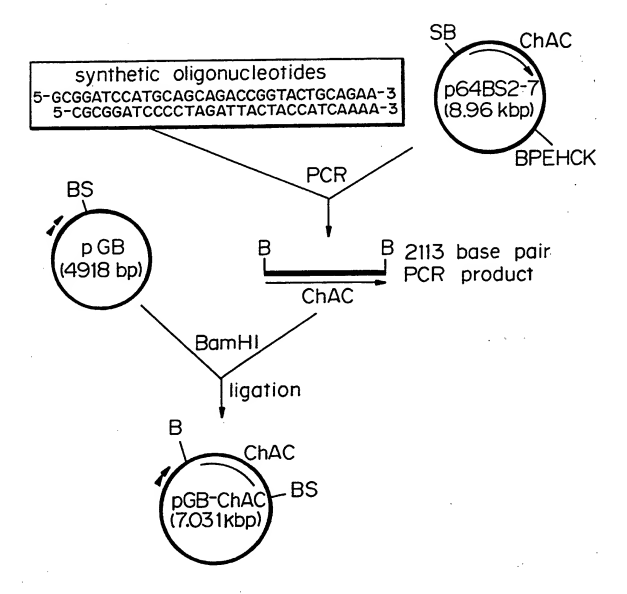


FIG. 2



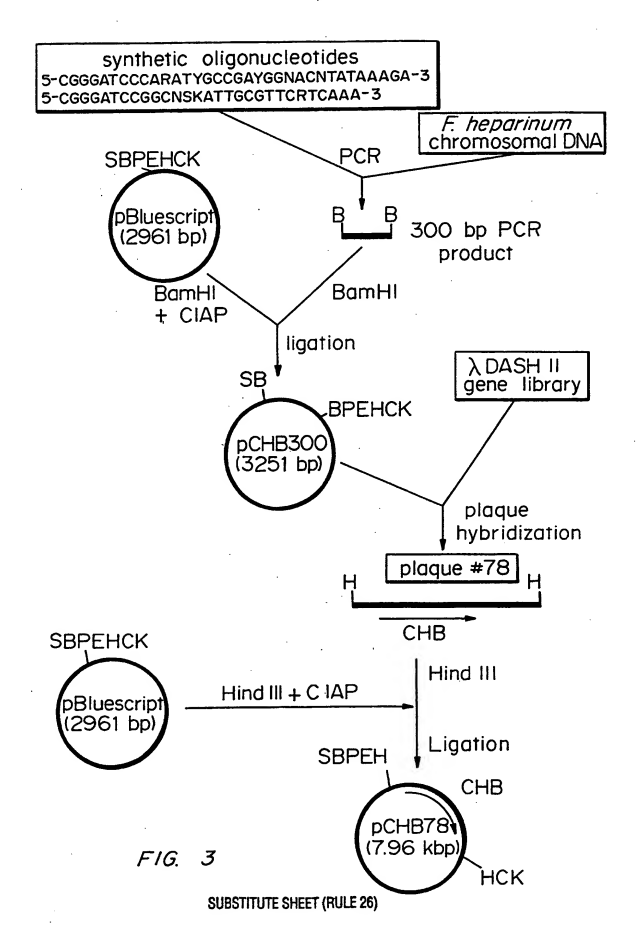
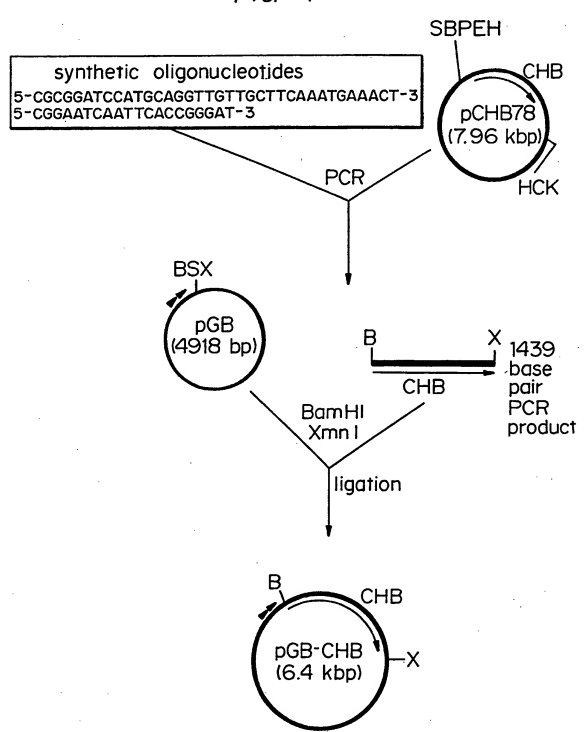


FIG. 4



### INTERNATIONAL SEARCH REPORT

Intermed Application No PCT/US 95/08560

CLASSIFICATION OF SUBJECT MATTER
PC 6 C12N9/88 C12N15/60 //(C12N9/88,C12R1:20) IPC 6 C12N9/88 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12N IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category \* BIOCHEM. J. (1975), 151(1), 121-9, MICHELACCI, YARA M. ET AL 'Comparative 1,2 X study between a chondroitinase B and a chondroitinase AC from Flavobacterium heparinium. Isolation of a chondroitinase AC-susceptible dodecasaccharide from chondroitin sulfate B' 3-19 see the whole document Y 3-18 DATABASE WPI Section Ch, Week 9419 Derwent Publications Ltd., London, GB; Class D16, AN 94-155922 & JP,A,06 098 769 ( TAIYO FISHERY CO LTD) , 12 April 1994 see abstract Patent family members are listed in annex. Further documents are listed in the continuation of box C. X "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means document published prior to the international filing date but later than the priority date claimed '&' document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 28. 11. 95 2 November 1995 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3016 Gurdjian, D

# INTERNATIONAL SEARCH REPORT

Int Smal Application No
PCT/US 95/08560

		PC1/US 95/08380	
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
ategory *	Citation of document, with indication, where appropriate, or the contract of		
•	DATABASE WPI Section Ch, Week 8046 Derwent Publications Ltd., London, GB; Class B04, AN 80-81758C & JP,A,55 127 988 ( SEIKAGAKU KK) , 4 October 1980 see abstract	19	
ſ	US,A,4 390 628 (JOHANSEN JACK T) 28 June 1983 see abstract	19	
X	BIOCHIM. BIOPHYS. ACTA (1987), 923(2), 291-301, MICHELACCI, YARA M. ET AL 'Isolation and characterization of an induced chondroitinase ABC from Flavobacterium heparinum' see the whole document	1,2	
		,	

2

# INTERNATIONAL SEARCH REPORT

Information on patent family members

Int mal Application No
PCT/US 95/08560

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
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		SU-A- 1184434	07-10-85
		US-A- 4388406	14-06-83
		AT-T- 5976	15-02-84
		EP-A,B 0019474	26-11-80
·	•	JP-C- 1586049	31-10-90
		JP-B- 2012556	20-03-90
		JP-A- 56035983	08-04-81
		JP-B- 1033158	12-07-89
		JP-C- 1549173	09-03-90
•		JP-A- 56035984	08-04-81
		US-A- 4340675	20-07-82